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RedMD – A New Package for Reduced Molecular Dynamics

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We developed a RedMD package to perform molecular dynamics simulations for coarse-grained models of proteins, nucleic acids and its complexes. Simulations can be carried out in the microcanonical ensemble, as well as with Berendsen and Langevin thermostats. We provide tools to generate initial configuration and topology which are based on the elastic network approach and its extensions. Topology generators can be modified by users to add for example a new potential type. The code is written in C/C++ languages and the structure/topology of a molecule is based on an XML format. The code is distributed under GNU public licence and will be available at <http://bionano.icm.edu.pl/>.

1 Introduction

We created an open-source, scalable package for reduced (coarse-grained) molecular dynamics (MD) simulations of biomolecules¹ on a micro- to mili-second time scales. It is written in C/C++ and parallelized with an OpenMP technology. To generate the topology and force field of a molecule we use an XML-based format. Currently implemented force field generators include the Elastic Network Model² and its anharmonic extensions for the ribosome³, nucleosome⁴ and HIV-1 protease⁵.

2 Molecular Dynamics

MD is a widely used technique to investigate the dynamical properties of molecules. It numerically solves in finite time steps the Newton's equations of motion and provides trajectories i.e., the coordinates and momenta of particles as a function of time. Our MD package generates coarse-grained representations of biopolymers in which entire groups of atoms are represented by single interacting centers (pseudo-atoms), (see Figure 1). In one bead models, one pseudo-atom can represent the whole amino acid or nucleotide. The coarse-graining procedure is applied to reduce the number of degrees of freedom, increase the integration time step, and achieve at least a microsecond MD simulation time scale. RedMD currently supports MD simulations in the microcanonical ensemble, in the canonical ensemble with the Berendsen thermostat, and with the Langevin bath thermal coupling. To enlarge the integration time step we implemented the SHAKE algorithm^{6,7} which was developed to satisfy the bond geometry constraints in MD simulations.

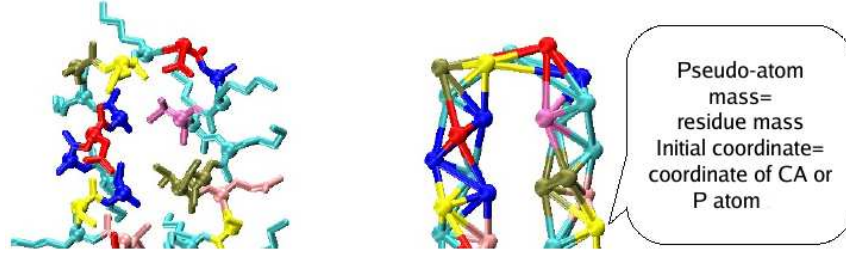


Figure 1. The coarse-graining procedure. Left: all-atom representation, right: reduced one-bead model.

2.1 Microcanonical Ensemble

In the microcanonical ensemble (NVE) we consider the Newton's equation:

$$\frac{d\vec{p}_i}{dt} = -\frac{\partial V}{\partial \vec{r}_i}, \quad i = 1, \dots, N$$

where V is a potential energy function which depends on nucleic coordinates r_i , p_i is the momentum of a particle i and N is the number of particles in the system. Numerical solution of this equation generates the trajectory of motion. We implemented two popular algorithms: velocity Verlet and Leap-Frog⁸ which are based on Taylor expansion.

2.2 Thermostats

To maintain constant temperature RedMD provides various thermostats. A common method of both thermal coupling and reproducing the contact with solvent molecules is Langevin Dynamics. We consider the equation:

$$\frac{d\vec{p}_i}{dt} = -\frac{\partial V}{\partial \vec{r}_i} - \gamma_i \vec{p}_i + \vec{R}(t), \quad i = 1, \dots, N$$

where γ_i is the collision parameter and $\vec{R}(t)$ is a random force vector satisfying:

$$\langle \vec{R}(t) \rangle = 0, \quad \langle \vec{R}(t) \vec{R}(t') \rangle = 2\gamma_i k_B T m_i \delta(t - t')$$

where k_B is the Boltzmann constant and T is the bath temperature. We solve this equation with Brünger-Brooks-Karplus integrator (BBK)⁹. Through the Langevin equation the system couples to a heat bath globally, but is also locally subjected to random noise. If we are interested in imposing the global coupling with minimal local disturbance the Langevin equation is modified to:

$$\frac{d\vec{p}_i}{dt} = -\frac{\partial V}{\partial \vec{r}_i} + m_i \gamma \left(\frac{T_0}{T} - 1 \right) \vec{p}_i, \quad i = 1, \dots, N$$

where T_0 is the bath temperature, T the current temperature and γ (in ps^{-1}) determines the strength of coupling with the thermal bath. This is known as the Berendsen thermostat¹⁰. The solution is based on scaling the momenta in each step by a factor

$$\lambda = \sqrt{1 + \frac{\Delta t}{\tau_T} \left(\frac{T_0}{T} - 1 \right)}$$

where τ_T is equal to $(2\gamma)^{-1}$.

2.3 Brownian Dynamics (BD)

RedMD provides a BD simulation based on the Ermak-McCammon algorithm¹¹. The trajectory is generated according to the equation:

$$\vec{r}_i(t + \Delta t) = \vec{r}_i(t) + \frac{D}{k_b T} \vec{F}_i(t) \Delta t + \vec{R}_i(t) \quad , \quad i = 1, \dots, N$$

where D is a diffusion coefficient of a molecule and $\vec{R}(t)$ is a random force vector satisfying:

$$\langle \vec{R}(t) \rangle = 0, \quad \langle \vec{R}(t) \vec{R}(t') \rangle = 6D\Delta t \delta(t - t').$$

3 Parallelization

Because the calculation of the non-bonded interactions and forces is the most time consuming step in MD, its optimization is very important. To parallelize this calculation we applied the OpenMP technology which uses a shared memory architecture. For the nucleosome model test force calculations for 1000 NVE simulation steps show that the speedup (defined as the ratio of time using one thread to the time of using N threads) for 7 cores is over 6 so it is almost linear.

4 The XML Force Field Format

The initial configuration and force field of the molecule are generated in an XML-based format. We provide utilities to produce the input XML file from the PDB or PDBML/XML files (<http://www.rcsb.org/pdb/>). Our XML format is a flexible way of representing the topology and force field. It contains information on atomic ids, masses, names, coordinates, and properties such as bonded and non-bonded potentials. It is possible to save any calculated values like temperature factors, forces or energies (see example below).

```
<?xml version="1.0"?>
<STRUCTURE>
<!-- Generated with: ./RedMD_genModel_Rib 1A36.xml -->
<NONBONDED>
  <NBMRSE cutoff="35.000000">
    <PAIR type1="CA" type2="CA" alpha="0.707000" E0="0.055055" l0="9.500000"/>
    <PAIR type1="P" type2="P" alpha="0.707000" E0="0.071348" l0="17.600000"/>
    <PAIR type1="CA" type2="P" alpha="0.707000" E0="0.062674" l0="12.930584"/>
  </NBMRSE>
</NONBONDED>
<MOLECULE molId="mol1">
  <GROUP label="basePairs" k="0.600000"/>
  <ATOM id="1" name="P" x="-50.7" y="76.7" z="327.1" resName="U" chainID="A" resSeq="2" m="305"/>
  <ATOM id="2" name="P" x="-50.8" y="73.7" z="332.1" resName="U" chainID="A" resSeq="3" m="305"/>
  <ATOM id="3" name="P" x="-48.5" y="75.0" z="336.9" resName="G" chainID="A" resSeq="4" m="344"/>
  <BOND idAtom1="1" idAtom2="2" k="3.0" l0="5.80"/>
  <BOND idAtom1="1" idAtom2="3" k="2.5" l0="10.15"/>
  <BOND idAtom1="1" idAtom2="4" k="0.5" l0="9.40"/>
  <MRSE idAtom1="1" idAtom2="5" alpha="0.707" l0="6.19" E0="0.71" mark="o:P:P"/>
  <MRSE idAtom1="1" idAtom2="6" alpha="0.707" l0="6.87" E0="0.63" mark="o:P:P"/>
  <MRSE idAtom1="1" idAtom2="7" alpha="0.707" l0="11.90" E0="0.27" mark="o:P:P"/>
</MOLECULE>
</STRUCTURE>
```

5 From PDB to Trajectory

First, one needs to convert a PDB or PDBML file to our XML file format with one of the programs distributed in the RedMD package. The user can choose from a few coarse-grained force fields. Second, the user needs to specify MD simulation parameters and then

can generate the trajectory. Currently supported trajectory output formats are XYZ, PDB, DCD and VEL (analogous to XYZ but saves velocities).

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